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ENTRY SESSION FULL ESTIMATED COST 0.45 0.66

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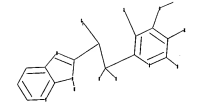
TSCA INFORMATION NOW CURRENT THROUGH December 2, 2006

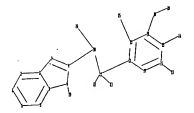
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http://www.cas.org/support/stngen/stndoc/properties.html

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chain nodes :
10 11 18 19 21 22 23 24 25 26 28
ring nodes :
1 2 3 4 5 6 7 8 9 12 13 14 15 16 17
chain bonds :
8-10 9-19 10-11 10-26 11-12 11-21 11-22 13-25 14-18 15-24 16-23 18-28
ring bonds :
1-2 1-6 2-3 3-4 3-7 4-5 4-9 5-6 7-8 8-9 12-13 12-17 13-14 14-15 15-16
16-17
exact/norm bonds :
3-7 4-9 7-8 8-9 10-26 14-18 18-28
exact bonds :
8-10 9-19 10-11 11-12 11-21 11-22 13-25 15-24 16-23
normalized bonds :
1-2 1-6 2-3 3-4 4-5 5-6 12-13 12-17 13-14 14-15 15-16 16-17
isolated ring systems :
containing 1 : 12 :
```

## G1:C,H

## Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:CLASS 11:CLASS 12:Atom 13:Atom 14:Atom 15:Atom 16:Atom 17:Atom 18:CLASS 19:CLASS 21:CLASS 22:CLASS 23:CLASS 24:CLASS 25:CLASS 26:CLASS 28:CLASS

=> d 11

L1 HAS NO ANSWERS

L1

Structure attributes must be viewed using STN Express query preparation.

=> s 11

SAMPLE SEARCH INITIATED 10:56:02 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED - 13 TO ITERATE

100.0% PROCESSED

13 ITERATIONS

5 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*

BATCH \*\*COMPLETE\*\*

PROJECTED ITERATIONS:

44 TO 476

PROJECTED ANSWERS:

5 TO 234

L2

5 SEA SSS SAM L1

=> s 11 full

FULL SEARCH INITIATED 10:56:07 FILE 'REGISTRY'

FULL SCREEN SEARCH COMPLETED - 256 TO ITERATE

100.0% PROCESSED

256 ITERATIONS

133 ANSWERS

SEARCH TIME: 00.00.01

L3 133 SEA SSS FUL L1

=> file caplus

COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION

FULL ESTIMATED COST

172.10 172.76

FILE 'CAPLUS' ENTERED AT 10:56:11 ON 02 MAY 2007

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http://www.cas.org/infopolicy.html

=> s 13 full

L4 7 L3

=> d ibib abs hitstr tot

L4 ANSWER 1 OF 7 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2006:341554 CAPLUS

DOCUMENT NUMBER: 144:381708

TITLE: In vivo characterization of the novel imidazopyridine

BYK191023 [2-[2-(4-methoxy-pyridin-2-yl)-ethyl]-3H-imidazo[4,5-b]pyridine], a potent and highly selective

inhibitor of inducible nitric-oxide synthase

AUTHOR(S): Lehner, Martin D.; Marx, Degenhard; Boer, Rainer;

Strub, Andreas; Hesslinger, Christian; Eltze, Manfrid; Ulrich, Wolf-Ruediger; Schwoebel, Frank; Schermuly,

Ralph Theo; Barsig, Johannes

CORPORATE SOURCE: Department of Pharmacology, ALTANA Pharma AG,

Konstanz, Germany

SOURCE: Journal of Pharmacology and Experimental Therapeutics

(2006), 317(1), 181-187

CODEN: JPETAB; ISSN: 0022-3565

PUBLISHER: American Society for Pharmacology and Experimental

Therapeutics

DOCUMENT TYPE: Journal LANGUAGE: English

Excessive release of nitric oxide from inducible nitric-oxide synthase (iNOS) has been postulated to contribute to pathol. in a number of inflammatory diseases. We recently identified imidazopyridine derivs. as a novel class of potent nitric-oxide synthase inhibitors with high selectivity for the inducible isoform. In the present study, we tested the in vivo potency of BYK191023 [2-[2-(4-methoxy-pyridin-2-yl)-ethyl]-3Himidazo[4,5-b]pyridine], a selected member of this inhibitor class, in three different rat models of lipopolysaccharide-induced systemic inflammation. Delayed administration of BYK191023 dose-dependently suppressed the lipopolysaccharide-induced increase in plasma nitrate/nitrite (NOx) levels with an ED50 of 14.9  $\mu$ mol/kg/h. model of systemic hypotension following high-dose lipopolysaccharide challenge, curative administration of BYK191023 at a dose that inhibited 83% of the NOx increase completely prevented the gradual decrease in mean arterial blood pressure observed in vehicle-treated control animals. vasopressor effect was specific for endotoxemic animals since BYK191023 did not affect blood pressure in saline-challenged controls. In addition, in a model of lipopolysaccharide-induced vascular hyporesponsiveness, BYK191023 infusion partially restored normal blood pressure responses to norepinephrine and sodium nitroprusside via an L-arginine competitive mechanism. Taken together, BYK191023 is a member of a novel class of highly isoform-selective iNOS inhibitors with promising in vivo activity suitable for mechanistic studies on the role of selective iNOS inhibition as well as clin. development.

RL: DMA (Drug mechanism of action); PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(in vivo characterization of inducible nitric-oxide synthase inhibitor imidazopyridine BYK191023)

RN 608880-48-4 CAPLUS

ΙT

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)ethyl]- (9CI) (CA INDEX NAME)

608880-48-4, BYK191023

REFERENCE COUNT: 39 THERE ARE 39 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 2 OF 7 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2006:43156 CAPLUS

DOCUMENT NUMBER: 144:163527

TITLE: The novel imidazopyridine 2-[2-(4-Methoxy-pyridin-2-

yl)-ethyl]-3H-imidazo[4,5-b]pyridine (BYK191023) is a

highly selective inhibitor of the inducible

nitric-oxide synthase

AUTHOR(S): Strub, Andreas; Ulrich, Wolf-Ruediger; Hesslinger,

Christian; Eltze, Manfrid; Fuchss, Thomas; Strassner,

Jochen; Strand, Susanne; Lehner, Martin D.; Boer,

Rainer

CORPORATE SOURCE: Departments of Biochemistry, Chemistry and

Pharmacology, ALTANA Pharma AG, Konstanz, Germany

SOURCE: Molecular Pharmacology (2006), 69(1), 328-337

CODEN: MOPMA3; ISSN: 0026-895X

PUBLISHER: American Society for Pharmacology and Experimental

Therapeutics

DOCUMENT TYPE: Journal LANGUAGE: English

AΒ We have identified imidazopyridine derivs. as a novel class of NO synthase inhibitors with high selectivity for the inducible isoform. 2-[2-(4-Methoxy-pyridin-2-yl)-ethyl]-3H-imidazo[4,5-b]pyridine (BYK191023) showed half-maximal inhibition of crudely purified human inducible (iNOS), neuronal (nNOS), and endothelial (eNOS) NO synthases at 86 nM, 17 μM, and 162  $\mu\text{M}\text{,}$  resp. Inhibition of inducible NO synthase was competitive with L-arginine, pointing to an interaction of BYK191023 with the catalytic center of the enzyme. In radioligand and surface plasmon resonance expts., BYK191023 exhibited an affinity for iNOS, nNOS, and eNOS of 450 nM, 30  $\mu$ M, and >500  $\mu$ M, resp. Inhibition of cellular nitrate/nitrite synthesis in RAW, rat mesangium, and human embryonic kidney 293 cells after iNOS induction showed 40- to 100-fold higher IC50 values than at the isolated enzyme, in agreement with the much higher L-arginine concns. in cell culture media and inside intact cells. BYK191023 did not show any toxicity in various rodent and human cell lines up to high micromolar concns. The inhibitory potency of BYK191023 was tested in isolated organ models of iNOS (lipopolysaccharide-treated and phenylephrine-precontracted rat aorta; IC50 =  $7 \mu M$ ), eNOS (arecaidine propargyl ester-induced relaxation of phenylephrine-precontracted rat aorta; IC50 > 100 µM), and nNOS (field-stimulated relaxation of

phenylephrine-precontracted rabbit corpus cavernosum; IC50 > 100  $\mu M)$ . These data confirm the high selectivity of BYK191023 for iNOS over eNOS and nNOS found at isolated enzymes. In summary, we have identified a new highly selective iNOS inhibitor structurally unrelated to known compds. and L-arginine. BYK191023 is a valuable tool for the investigation of iNOS-mediated effects in vitro and in vivo.

IT 608880-48-4, BYK 191023

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(structure activity relationship studied of imidazopyridine compds. as selective inhibitors of nitric-oxide synthase isoforms)

RN 608880-48-4 CAPLUS

CN

1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)ethyl]- (9CI) (CA INDEX NAME)

REFERENCE COUNT: 40 THERE ARE 40 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 3 OF 7 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2005:300447 CAPLUS

DOCUMENT NUMBER: 142:373838

TITLE: Preparation of imidazopyridine derivatives as

inducible NO-synthase inhibitors

INVENTOR(S): Fuchss, Thomas; Martin, Thomas; Boer, Rainer; Strub,

Andreas; Eltze, Manfrid; Lehner, Martin; Ulrich,

Wolf-Ruediger

PATENT ASSIGNEE(S): Altana Pharma A.-G., Germany

SOURCE: PCT Int. Appl., 34 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.				KIND DATE					APPI	ICAT	ION :		DATE						
WO 2005030771			A1 20050407					WO 2	004-	EP52		20040930							
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		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,		
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	KZ,	LC,		
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,	NI,		
							•				SC,								
		ТJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW		
	RW:	BW,	GH,	GM,	ΚE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,		
											BE,								
											LU,								
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	2007										006-								
	2006								NO 2006-1344 2006										
US	2007	0430	/3		A1		2007	0222	US 2006-573484 20060324							324			

IN 2006MN00475
PRIORITY APPLN. INFO.:

20070316

IN 2006-MN475 EP 2003-22053 20060424 A 20031001

20040930

OTHER SOURCE(S):

WO 2004-EP52378 W

GI

RN

CASREACT 142:373838; MARPAT 142:373838

Ι

AB Title compds. I [R1 = H, alkyl; R2 = H, alkyl; R3 = H, halo; R4 = H, halo, alkyl, alkoxy; R5 = alkyl; A = alkylene] and their resp. pharmaceutically acceptable salts, are prepared and disclosed as inducible no-synthase inhibitors. Thus, e.g., II was prepared via Suzuki coupling of 2-[2-(4-methoxypyridin-2-yl)ethyl]-6-iodo-3H-imidazo[4,5-b]pyridine (preparation given) with N,N-dimethyl-4-bromobenzenesulfonamide. The activity of I towards inducible NO-synthase was evaluated in inhibition assays and revealed -logIC50 values in the range of 7.45 up to 7.86 mol/L. I as inducible NO-synthase inhibitors should prove useful in the treatment of acute and chronic inflammatory diseases.

II

IT 849357-47-7P 849357-48-8P 849357-49-9P 849357-50-2P 849357-51-3P 849357-52-4P 849357-54-6P 849357-55-7P 849357-56-8P 849357-57-9P 849357-58-0P 849357-59-1P RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of imidazopyridine derivs. as inducible NO-synthase inhibitors) 849357-47-7 CAPLUS

CN Benzenesulfonamide, 4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]-N,N-dimethyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c}
N & N & CH_2 - CH_2 \\
\hline
NH & NH & N
\end{array}$$

RN 849357-48-8 CAPLUS

CN Benzenesulfonamide, N, N-diethyl-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c}
N & N \\
N & N$$

RN 849357-49-9 CAPLUS

CN Benzenesulfonamide, 4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]-N-methyl- (9CI) (CA INDEX NAME)

RN 849357-50-2 CAPLUS

CN Benzenesulfonamide, 4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ \end{array}$$

RN 849357-51-3 CAPLUS

CN Benzenesulfonamide, N-ethyl-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]- (9CI) (CA INDEX NAME)

RN 849357-52-4 CAPLUS

CN Benzenesulfonamide, 2-fluoro-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]-N,N-dimethyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} O & F \\ Me_2N - S & H \\ O & N & N \end{array}$$
 OME

RN 849357-54-6 CAPLUS

CN Benzenesulfonamide, 2-fluoro-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]- (9CI) (CA INDEX NAME)

RN 849357-55-7 CAPLUS

CN Benzenesulfonamide, 2-fluoro-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]-N-methyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} O & F \\ \hline MeNH-S & H \\ O & N \end{array} \quad CH_2-CH_2 \quad OMe$$

RN '849357-56-8 CAPLUS

CN Benzenesulfonamide, N-ethyl-2-fluoro-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} C & F \\ C & N \end{array}$$

RN 849357-57-9 CAPLUS

CN Benzenesulfonamide, N-ethyl-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]-N-methyl- (9CI) (CA INDEX NAME)

RN 849357-58-0 CAPLUS

CN Benzenesulfonamide, N,N-diethyl-2-fluoro-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} C & F & H & CH_2 - CH_2 & OMe \\ \hline & N & N & N & N \end{array}$$

RN 849357-59-1 CAPLUS

CN Benzenesulfonamide, N-ethyl-2-fluoro-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]-N-methyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} \text{Me O} & \text{F} \\ & \parallel & \parallel \\ \text{Et-N-S} \\ & \parallel & \parallel \\ & \text{O} \end{array}$$

IT 608880-54-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of imidazopyridine derivs. as inducible NO-synthase inhibitors)

RN 608880-54-2 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 6-iodo-2-[2-(4-methoxy-2-pyridinyl)ethyl]-(9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 4 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 4 OF 7 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2005:300446 CAPLUS

DOCUMENT NUMBER:

142:373837

TITLE:

Preparation of imidazopyridine derivatives as

inducible NO-synthase inhibitors

INVENTOR(S):

Fuchss, Thomas; Martin, Thomas; Boer, Rainer; Strub, Andreas; Eltze, Manfrid; Lehner, Martin; Ulrich,

Wolf-Ruediger

PATENT ASSIGNEE(S):

Altana Pharma A.-G., Germany

SOURCE:

PCT Int. Appl., 66 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA	TENT	NO.			KIND DATE				APPL	ICAT	ION I		DATE							
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								PT,												
								UA,												
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								ТJ,												
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									CA 2004-2540243 EP 2004-787262											
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NO 2006001317					ν Τ															
	US 2006293302 ORITY APPLN. INFO.:						2000	1220												
101(11	ORIII APPLIN. INFO.:									EP 2003-22046										
HER S	IER SOURCE(S):					WO 2004-EP52377 W 20040930 CASREACT 142:373837; MARPAT 142:373837														

GI

AB Title compds. I [R1 = H, alkyl, cycloalkyl, etc.; R2 = H, alkoxyalkyl, hydroxyalkyl, etc.; R3 = alkyl, CF3, completely or predominantly F-substituted alkoxy, etc.; R1 and R2 together = (un)saturated-, (un)substituted-nitrogen heterocycle; R4 = H, halo, alkyl, alkoxy; R5 = alkyl; A = alkylene] and their resp. pharmaceutically acceptable salts, are prepared and disclosed as inducible no-synthase inhibitors. Thus, e.g., II was prepared via Suzuki coupling of 2-[2-(4-methoxypyridin-2-yl)ethyl]-6-iodo-3H-imidazo[4,5-b]pyridine (preparation given) with 1-(4-bromo-benzene-sulfonyl)-4-methyl-piperazine. The activity of I towards inducible NO-synthase was evaluated in inhibition assays and revealed -logIC50 values in the range of 6.51 up to 7.89 mol/L. I as inducible NO-synthase inhibitors should prove useful in the treatment of acute and chronic inflammatory diseases.

Ι

IT849530-66-1P 849530-68-3P 849530-70-7P 849530-72-9P 849530-74-1P 849530-76-3P 849530-78-5P 849530-80-9P 849530-82-1P 849530-84-3P 849530-86-5P 849530-88-7P 849530-90-1P 849530-92-3P 849530-94-5P 849530-96-7P 849530-98-9P 849531-00-6P 849531-02-8P 849531-04-0P 849531-06-2P 849531-08-4P 849531-10-8P 849531-12-0P 849531-14-2P 849531-16-4P 849531-18-6P 849531-20-0P 849531-23-3P 849531-25-5P 849531-27-7P 849531-29-9P 849531-31-3P 849531-33-5P 849531-36-8P 849531-38-0P 849531-40-4P 849531-42-6P 849531-44-8P 849531-46-0P 849531-48-2P 849531-50-6P 849531-52-8P 849531-54-0P 849531-56-2P 849531-58-4P 849531-60-8P 849531-62-0P 849531-64-2P 849531-66-4P 849531-68-6P 849531-70-0P 849531-72-2P 849531-74-4P 849531-76-6P 849531-78-8P 849531-80-2P 849531-82-4P 849531-84-6P 849531-86-8P 849531-88-0P 849531-90-4P 849531-92-6P 849531-94-8P 849531-96-0P 849531-98-2P 849532-00-9P RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of imidazopyridine derivs. as inducible NO-synthase inhibitors)

RN 849530-66-1 CAPLUS

CN Piperazine, 1-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]-4-methyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ N & & S \\ \hline & & \\ N & & \\ \end{array}$$

RN 849530-68-3 CAPLUS

CN Piperazine, 1-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]-4-(phenylmethyl)- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ &$$

RN 849530-70-7 CAPLUS

CN Piperazine, 1-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]-4-phenyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} O & H & CH_2-CH_2 \\ \hline \\ Ph & N \end{array}$$

RN 849530-72-9 CAPLUS

CN Piperazine, 1-(4-cyanophenyl)-4-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

RN 849530-74-1 CAPLUS

CN Piperazine, 1-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]-4-(4-methylphenyl)- (9CI) (CA INDEX NAME)

RN 849530-76-3 CAPLUS

CN Piperazine, 1-(2,4-dimethylphenyl)-4-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

RN 849530-78-5 CAPLUS

CN Piperazine, 1-(3,5-dichlorophenyl)-4-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} C1 & O \\ N & S \\ O & NH \end{array}$$

RN 849530-80-9 CAPLUS

CN Piperazine, 1-(2-methoxyethyl)-4-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ \text{MeO-CH}_2\text{-CH}_2 \end{array} \\ & & & \\ & & & \\ \text{OMe} \\ & & & \\ \end{array}$$

RN . 849530-82-1 CAPLUS

CN Piperazine, 1-acetyl-4-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

Ac 
$$N - S - CH_2 - CH_2$$
 OMe

RN 849530-84-3 CAPLUS

CN Morpholine, 4-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} O & & & \\ N & S & & \\ O & & & \\ N & & \\ \end{array}$$

RN 849530-86-5 CAPLUS

CN 1H-1,4-Diazepine, hexahydro-1-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]-4-methyl- (9CI) (CA INDEX NAME)

RN 849530-88-7 CAPLUS

CN Piperidine, 1-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]-4-methyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ N & & \\ \end{array}$$

RN 849530-90-1 CAPLUS

CN Piperidine, 4-benzoyl-1-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

RN 849530-92-3 CAPLUS

CN 1,4-Dioxa-8-azaspiro[4.5]decane, 8-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

RN 849530-94-5 CAPLUS

CN Isoquinoline, 1,2,3,4-tetrahydro-6,7-dimethoxy-2-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

RN 849530-96-7 CAPLUS

CN 5H-1,4-Diazepin-5-one, hexahydro-1-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

RN 849530-98-9 CAPLUS

CN Benzenesulfonamide, N-(2-hydroxyethyl)-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]- (9CI) (CA INDEX NAME)

RN 849531-00-6 CAPLUS

CN Benzenesulfonamide, N,N-bis(2-hydroxyethyl)-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]- (9CI) (CA INDEX NAME)

RN 849531-02-8 CAPLUS

CN Benzenesulfonamide, 4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]-N-(phenylmethyl)- (9CI) (CA INDEX NAME)

RN 849531-04-0 CAPLUS

CN Benzenesulfonamide, N-cyclohexyl-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]- (9CI) (CA INDEX NAME)

RN 849531-06-2 CAPLUS

CN Benzenesulfonamide, 4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]-N,N-dimethyl-2-(trifluoromethoxy)- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ &$$

RN 849531-08-4 CAPLUS

CN Benzenesulfonamide, 4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]-N,N-dimethyl-2-(trifluoromethyl)- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c}
 & H \\
 & N \\
 & N \\
 & N \\
 & O \\$$

RN 849531-10-8 CAPLUS

CN Benzenesulfonamide, 4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]-N,N,3-trimethyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} Me \\ \hline \\ Me_2N-S \\ \hline \\ O \end{array}$$

RN 849531-12-0 CAPLUS

CN Benzenesulfonamide, 4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]-N-phenyl- (9CI) (CA INDEX NAME)

RN 849531-14-2 CAPLUS

CN Benzenesulfonamide, 4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]-N-(4-methylphenyl)- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} \text{Me} & \text{O} & \text{H} \\ \text{NH} & \text{S} \\ \text{O} & \text{N} \\ \text{N} & \text{N} \end{array}$$

RN 849531-16-4 CAPLUS

CN Benzenesulfonamide, N-(2-methoxyphenyl)-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo(4,5-b)pyridin-6-yl]- (9CI) (CA INDEX NAME)

RN 849531-18-6 CAPLUS

CN Benzenesulfonamide, N-[4-(dimethylamino)phenyl]-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} \text{Me}_2\text{N} & \text{O} \\ \text{NH-S} & \text{H} \\ \text{O} & \text{NH-CH}_2\text{-CH}_2 \end{array}$$
 OMe

RN 849531-20-0 CAPLUS

CN Benzenesulfonamide, N-(4-chlorophenyl)-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]-N-methyl- (9CI) (CA INDEX NAME)

RN 849531-23-3 CAPLUS

CN Piperazine, 1-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]-4-(2-phenylethyl)- (9CI) (CA INDEX NAME)

RN 849531-25-5 CAPLUS

CN Piperazine, 1-ethyl-4-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c}
 & O \\
 & N \\
 & N \\
 & O \\
 & N \\
 & N \\
 & N \\
 & O \\$$

RN 849531-27-7 CAPLUS

CN Piperazine, 1-(2,6-dimethylphenyl)-4-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

RN 849531-29-9 CAPLUS

CN Piperazine, 1-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]-4-(2-methylphenyl)- (9CI) (CA INDEX NAME)

RN 849531-31-3 CAPLUS

CN Piperazine, 1-[[3-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]-4-methyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ N & & \\ N & & \\ \end{array}$$

RN 849531-33-5 CAPLUS

CN Piperidine, 1-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c}
 & O \\
 & N \\
 & S \\
 & O \\
 & O \\
 & N \\
 & N \\
 & O \\$$

RN 849531-36-8 CAPLUS

CN Piperazine, 1-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]-2-(trifluoromethoxy)phenyl]sulfonyl]-4-methyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c}
 & O & O - CF3 \\
 & N & S & H \\
 & N & N & CH_2 - CH_2 & OMe
\end{array}$$

RN 849531-38-0 CAPLUS

CN Isoquinoline, 6,7-diethoxy-1,2,3,4-tetrahydro-2-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

EtO 
$$N = S$$
  $N = S$   $N = S$ 

RN 849531-40-4 CAPLUS

CN Piperazine, 1-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]-2-(trifluoromethyl)phenyl]sulfonyl]-4-methyl- (9CI) (CA INDEX NAME)

RN 849531-42-6 CAPLUS

CN Piperazine, 1-[[2-fluoro-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]-4-methyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ &$$

RN 849531-44-8 CAPLUS

CN Piperazine, 1-[[2-chloro-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]-4-methyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c}
 & O \\
 & N \\
 & N \\
 & O \\$$

RN 849531-46-0 CAPLUS

CN Piperazine, 1-[[3-fluoro-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]-4-methyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & \\ N & & \\ N & & \\ \end{array}$$

RN 849531-48-2 CAPLUS

CN 5H-1,4-Diazepin-5-one, hexahydro-1-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]-4-(phenylmethyl)- (9CI) (CA INDEX NAME)

RN 849531-50-6 CAPLUS

CN Benzenesulfonamide, 4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]-N-methyl-N-phenyl- (9CI) (CA INDEX NAME)

RN 849531-52-8 CAPLUS

CN Piperazine, 1-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]-3-methylphenyl]sulfonyl]-4-methyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ &$$

RN 849531-54-0 CAPLUS

CN 5H-1,4-Diazepin-5-one, hexahydro-1-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]-4-methyl- (9CI) (CA INDEX NAME)

Me N 
$$N = S$$
  $N = S$   $N = S$ 

RN 849531-56-2 CAPLUS

CN 5H-1,4-Diazepin-5-one, 4-ethylhexahydro-1-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

RN 849531-58-4 CAPLUS

CN Benzenesulfonamide, 4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]-N-(2-methylphenyl)- (9CI) (CA INDEX NAME)

RN 849531-60-8 CAPLUS

CN Benzenesulfonamide, 4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]-N-methyl-N-4-pyridinyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} Me & O \\ \hline \\ N & S \\ \hline \\ N & O \\ \end{array}$$

RN 849531-62-0 CAPLUS

CN Benzenesulfonamide, 4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]-N-methyl-N-(4-methylphenyl)- (9CI) (CA INDEX NAME)

Me O 
$$H$$
  $CH_2-CH_2$  OMe

RN 849531-64-2 CAPLUS

CN Benzenesulfonamide, N-[4-(dimethylamino)phenyl]-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]-N-methyl- (9CI) (CA INDEX NAME)

RN 849531-66-4 CAPLUS

CN Benzenesulfonamide, N-(2-fluoro-4-methylphenyl)-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ &$$

RN 849531-68-6 CAPLUS

CN Benzenesulfonamide, N-(4-methoxyphenyl)-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]- (9CI) (CA INDEX NAME)

RN 849531-70-0 CAPLUS

CN Benzenesulfonamide, N-(4-methoxyphenyl)-4-[2-[2-(4-methoxy-2pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]-N-methyl- (9CI) (CA INDEX NAME)

849531-72-2 CAPLUS RN

Benzenesulfonamide, 4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-CN b]pyridin-6-yl]-N-methyl-N-(2-methylphenyl)- (9CI) (CA INDEX NAME)

RN849531-74-4 CAPLUS

CN Benzenesulfonamide, N-(4-chlorophenyl)-4-[2-[2-(4-methoxy-2pyridinyl)ethyl]-1H-imidazo(4,5-b)pyridin-6-yl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} C1 & O & H & N \\ \hline & NH - S & N \\ \hline & O & N \\ \hline & N \\ & N \\ & N \\ \end{array}$$
 CH<sub>2</sub>- CH<sub>2</sub>- CH<sub>2</sub>- OMe

RN

849531-76-6 CAPLUS Pyrrolidine, 1-[[4-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-methoxy-2-pyridinyl)ethyl]CN b]pyridin-6-yl]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c}
 & O \\
 & N \\
 & S \\
 & O \\
 & N \\
 & N \\
 & N \\
 & N \\
 & O \\$$

RN 849531-78-8 CAPLUS

CN Azetidine, 1-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

RN 849531-80-2 CAPLUS

CN Benzenesulfonamide, N,N-bis(2-methoxyethyl)-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]- (9CI) (CA INDEX NAME)

RN 849531-82-4 CAPLUS

CN Benzenesulfonamide, N-cyclobutyl-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]- (9CI) (CA INDEX NAME)

RN 849531-84-6 CAPLUS

CN Benzenesulfonamide, N-cyclopropyl-4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} O & H & N \\ \hline O & NH - S & N \\ \hline O & N & N \\ \hline \end{array}$$

RN 849531-86-8 CAPLUS

CN Pyrrolidine, 1-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-7-methyl-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

RN 849531-88-0 CAPLUS

CN Piperidine, 1-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-7-methyl-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} O & Me \\ N & S \\ O & N \\ N & N \end{array}$$

$$CH_2 - CH_2$$

$$OMe$$

RN 849531-90-4 CAPLUS

CN Morpholine, 4-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-7-methyl-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} O & Me \\ N & S \\ \hline O & N \\$$

RN 849531-92-6 CAPLUS

CN Azetidine, 1-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-7-methyl-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

$$O = S$$

$$Me$$

$$N$$

$$N$$

$$CH_2 - CH_2$$

$$OMe$$

RN 849531-94-8 CAPLUS

CN Thiomorpholine, 4-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} O & H & N \\ \hline S & O & H \\ \hline S & N & N \\ \hline \end{array}$$
 
$$\begin{array}{c|c} CH_2 - CH_2 & OMe \\ \hline \end{array}$$

RN 849531-96-0 CAPLUS

CN Thiomorpholine, 4-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]-, 1-oxide (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

RN 849531-98-2 CAPLUS

CN Thiomorpholine, 4-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]-, 1,1-dioxide (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} O & & & \\ \hline O & & \\ O$$

RN 849532-00-9 CAPLUS

CN Isoquinoline, 1,2,3,4-tetrahydro-2-[[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} O & \\ \hline \\ N & \\ O & \\ \hline \\ N & \\ N & \\ \end{array}$$

IT 608880-54-2P, 2-[2-[4-Methoxypyridin-2-yl]ethyl]-6-iodo-3H-

imidazo[4,5-b]pyridine

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of imidazopyridine derivs. as inducible NO-synthase inhibitors)

RN 608880-54-2 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 6-iodo-2-[2-(4-methoxy-2-pyridinyl)ethyl]-(9CI) (CA INDEX NAME)

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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ACCESSION NUMBER: 2005:300445 CAPLUS

DOCUMENT NUMBER: 142:373836

TITLE: Preparation of imidazopyridine derivatives as

inducible NO-synthase inhibitors

INVENTOR(S): Fuchss, Thomas; Martin, Thomas; Boer, Rainer; Strub,

Andreas; Eltze, Manfrid; Lehner, Martin; Marx,

Degenhard; Ulrich, Wolf-Ruediger

PATENT ASSIGNEE(S): Altana Pharma A.-G., Germany

SOURCE: PCT Int. Appl., 41 pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA'	rent :	NO.			KIND DATE					APPL	ICAT	ION 1		DATE						
WO	2005	A1 20050407				WO 2	004-	EP52	20040930											
	W:	AE,	AG,	AL,	AM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,			
		CN,	co,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,			
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,			
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CN	CN 1856494													EE, HU, PL, SK, HR 20040930						
	BR 2004014873													20040930						
JP	JP 2007507465						2007	0329												
	NO 2006001316																			
	IN 2006MN00473						2007													
	RIORITY APPLN. INFO.:										003-									
			•	• •							004-					0040				
OTHER SO	THER SOURCE(S):						142.	3738		2	J J J		20040550							

OTHER SOURCE(S): MARPAT 142:373836

GI

Title compds. I [R1 = alkoxy; A = alkylene; R2 = H, halo, alkyl, alkoxy; Het = (un)substituted monocyclic or fused 5-10 membered (un)saturated heteroaryl containing 1-3 heteroatoms selected from N, O, and S] and their resp. pharmaceutically acceptable salts, are prepared and disclosed as inducible no-synthase inhibitors. Thus, e.g., II was prepared via Suzuki coupling of 2-[2-(4-methoxypyridin-2-yl)ethyl]-6-iodo-3H-imidazo[4,5-b]pyridine (preparation given) with 2-furanylboronic acid. The activity of I towards inducible NO-synthase was evaluated in inhibition assays and revealed -logIC50 values from 6.61 up to 7.61 mol/L. I as inducible NO-synthase inhibitors should prove useful in the treatment of acute and chronic inflammatory diseases.

IT 849356-71-4P 849356-72-5P 849356-73-6P 849356-74-7P 849356-75-8P 849356-76-9P 849356-77-0P 849356-78-1P 849356-79-2P 849356-80-5P 849356-81-6P 849356-82-7P 849356-83-8P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of imidazopyridine derivs. as inducible NO-synthase inhibitors) 849356-71-4 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 6-(2-furanyl)-2-[2-(4-methoxy-2-pyridinyl)ethyl]- (9CI) (CA INDEX NAME)

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$$\begin{array}{c|c} O & H \\ \hline & N \\ \hline & N \\ \end{array} CH_2 - CH_2 \\ \hline & N \\ \end{array} OMe$$

RN

RN 849356-72-5 CAPLUS
CN 1H-Imidazo[4,5-b]pyridine, 6-(3-furanyl)-2-[2-(4-methoxy-2-pyridinyl)ethyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & H & CH_2 - CH_2 \\ \hline & N & N \end{array}$$

RN 849356-73-6 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)ethyl]-6-(4-pyridinyl)- (9CI) (CA INDEX NAME)

RN 849356-74-7 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)ethyl]-6-(2-thienyl)- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} S & H & CH_2 - CH_2 \\ \hline & N & \cdot & N \end{array}$$

RN 849356-75-8 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)ethyl]-6-(3-thienyl)- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & H & CH_2-CH_2 & OMe \\ \hline & N & N & N & \end{array}$$

RN 849356-76-9 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 6-(1H-indol-5-yl)-2-[2-(4-methoxy-2-pyridinyl)ethyl]- (9CI) (CA INDEX NAME)

RN 849356-77-0 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 6-(1H-benzimidazol-2-yl)-2-[2-(4-methoxy-2-pyridinyl)ethyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & N & \\ & & CH_2-CH_2 \\ \end{array}$$

RN 849356-78-1 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridiny1)ethy1]-6-(5-methyl-1H-benzimidazol-2-yl)- (9CI) (CA INDEX NAME)

RN 849356-79-2 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 6-benzo[b]thien-3-yl-2-[2-(4-methoxy-2-pyridinyl)ethyl]- (9CI) (CA INDEX NAME)

RN 849356-80-5 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 6-(2-benzofuranyl)-2-[2-(4-methoxy-2-pyridinyl)ethyl]- (9CI) (CA INDEX NAME)

RN 849356-81-6 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 6-benzo[b]thien-2-yl-2-[2-(4-methoxy-2-pyridinyl)ethyl]- (9CI) (CA INDEX NAME)

RN 849356-82-7 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)ethyl]-6-(5-methyl-2-thienyl)- (9CI) (CA INDEX NAME)

Me 
$$S$$
  $N$   $CH_2-CH_2$   $N$   $N$   $N$   $N$ 

RN 849356-83-8 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)ethyl]-6-(5-phenyl-2-thienyl)- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} \text{Ph} & \text{S} & \text{H} \\ \hline & \text{N} & \text{CH}_2 - \text{CH}_2 \\ \hline & \text{N} & \text{N} \end{array}$$

IT 608880-54-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of imidazopyridine derivs. as inducible NO-synthase inhibitors)

RN 608880-54-2 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 6-iodo-2-[2-(4-methoxy-2-pyridinyl)ethyl]-(9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 6 OF 7 CAPLUS COPYRIGHT 2007 ACS on STN

3

ACCESSION NUMBER:

2005:300444 CAPLUS

DOCUMENT NUMBER:

142:373835

TITLE:

Preparation of imidazopyridine derivatives as

inducible NO-synthase inhibitors

INVENTOR(S):

Boer, Rainer; Ulrich, Wolf-Ruediger; Eltze, Manfrid;

Marx, Degenhard; Graedler, Ulrich; Fuchss, Thomas

PATENT ASSIGNEE(S):

Altana Pharma A.-G., Germany

SOURCE:

PCT Int. Appl., 31 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

1

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE \_\_\_\_ WO 2005030768 A1 20050407 WO 2004-EP52370 20040930 AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE,

SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG AU 2004276012 Α1 20050407 AU 2004-276012 20040930 CA 2540239 Α1 20050407 CA 2004-2540239 20040930 EP 1675853 Α1 20060705 EP 2004-787257 20040930 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, PL, SK, CN 1856492 Α 20061101 CN 2004-80027796 20040930 BR 2004015038 Α 20061212 BR 2004-15038 20040930 JP 2007507463 Т 20070329 JP 2006-530260 20040930 NO 2006001343 Α 20060324 NO 2006-1343 · 20060324 US 2007010549 Α1 20070111 US 2006-573203 20060324 PRIORITY APPLN. INFO.: EP 2003-22042 20031001 Α WO 2004-EP52370 W 20040930 OTHER SOURCE(S): MARPAT 142:373835

$$R^2$$
 $N$ 
 $R^1$ 
 $R^5$ 
 $R^3$ 
 $N$ 
 $N$ 
 $N$ 
 $N$ 
 $N$ 

GI

AB Title compds. I [R1 = H, alkyl; R2 = H, halo, OH, etc.; R3 = H, halo, alkyl, alkoxy; R4 = alkyl; R5 = alkyl] and their resp. pharmaceutically acceptable salts, are prepared and disclosed as inducible no-synthase inhibitors. Thus, e.g., II was prepared via Wittig reaction of triphenyl-{1-(3H-imidazo[4,5-b]pyridin-2-yl)-propyl}-phosphonium chloride (preparation given) with 4-methoxypyridine-2-carboxaldehyde and subsequent hydrogenation. The activity of II towards inducible NO-synthase was evaluated in an inhibition assay and revealed a -logIC50 value of 7.15 mol/L. I as inducible NO-synthase inhibitors should prove useful in the treatment of acute and chronic inflammatory diseases.

Ι

849346-44-7P 849346-45-8P 849346-46-9P RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of imidazopyridine derivs. as inducible NO-synthase inhibitors) RN 849346-44-7 CAPLUS

•x HCl

RN 849346-45-8 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[1-[(4-methoxy-2-pyridinyl)methyl]propyl]-(9CI) (CA INDEX NAME)

RN 849346-46-9 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)-1,1-dimethylethyl]-(9CI) (CA INDEX NAME)

IT 849346-55-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of imidazopyridine derivs. as inducible NO-synthase inhibitors)

RN 849346-55-0 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)-1-methylethyl]-(9CI) (CA INDEX NAME)

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 7 OF 7 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2003:777790 CAPLUS

DOCUMENT NUMBER: 139:292156

TITLE: Preparation of alkoxypyridines as inducible nitric

oxide synthase (iNOS) inhibitors

INVENTOR(S): Boer, Rainer; Marx, Degenhard; Eltze, Manfrid; Klein,

Thomas; Nave, Ruediger; Graedler, Ulrich; Fuchss, Thomas; Barsig, Johannes; Ulrich, Wolf-Ruediger

PATENT ASSIGNEE(S): Altana Pharma A.-G., Germany

SOURCE:

PCT Int. Appl., 52 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATI	ENT N	KINI	)	DATE		APP	LICAT		DATÉ										
WO 2	2003080607						2003	1002		WO	2003-	EP30	20030325						
								, DZ,											
	IS, JP, KR,					LV,	MA,	MK,	MX,	NO	, NZ,	PH,	PL,	SG,	TN,	UA,	US,		
		•		ZA,															
											, AT,								
						FR,	GB,	GR,	HU,	11	, IT,	LU,	MC,	NL,	PT,	· RO,	SE,		
an a				TR			2002	1000		~~	0000	0400	0000000						
CA A	24803	85			AI		2003	1002		CA	2003~	2480	20030325						
	U 2003226706					A1 20031008					2003-	2267	20030325						
EP :													20030325						
											, IT,								
						FI,	RO,	MK,	CY,	AL	, TR,	BG,	CZ,	EE,	HU,	SK			
BR 2	20030	0878	35		Α	2005	0111		BR	2003-	8785			2	0030				
CN :	16429.	55			Α	2005	0720		CN	2003-	8069	20030325							
US 2	20051	7112	25		A1							20030325							
JP 2	20055	2538	38		T		2005							20030325					
	53595				Α		2006	0526						20030325					
IN 2	2004MI	N004	162		Α		2005	20050218							20040820				
	71383				B2		2006	1121		US	2004-	5093	96		2004092				
NO 2	NO 2004004633						2004	1223		NO	2004-	4633			2	20040324			
	PRIORITY APPLN. INFO.:								NO 2004-4633 EP 2002-7049										
											2003-					0030			
OTHER SOU	OTHER SOURCE(S):					PAT	139:	29215	56				-						

Title compds. I [wherein R1 = alkoxy; A = alkylene; B = (un)substituted 3H-imidazo[4,5-b]pyridin-2-yl, 9H-purin-8-yl; their salts, N-oxides, and AB salts of the N-oxides] were prepared as inducible NO-synthase (iNOS) inhibitor for treatment of acute inflammatory diseases and chronic inflammatory diseases of peripheral organs and central nervous system (CNS). For example, II (m.p. =  $116-117^{\circ}$ ) was prepared by cyclocondensation of Me 3-(4-methoxypyridin-2-yl) propionate (preparation given) with 2,3-diaminopyridine in the presence of polyphosphoric acid at

160° for 1 h. Selected invention compds. inhibited iNOS with -logIC50 (M) in the range of 7.03-7.55. Thus, I and their pharmaceutical compns. are useful for treating acute inflammatory diseases, chronic inflammatory diseases of peripheral organs and CNS and cancer (no data).

ΙT 608880-48-4P, 2-[2-(4-Methoxypyridin-2-y1)ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1)ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1)ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1)ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1)ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1)ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1)ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1)ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1)ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1)ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1)ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1)ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1]ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1]ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1]ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1]ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1]ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1]ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1]ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1]ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1]ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1]ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1]ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1]ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1]ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1]ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1]ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1]ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1]ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1]ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1]ethy1]ethy1]-3H-imidazo[4,5-methoxypyridin-2-y1]ethy1]etb]pyridine 608880-53-1P, 2-[2-(4-Methoxypyridin-2-yl)ethyl]-6bromo-3H-imidazo[4,5-b]pyridine 608880-54-2P, 2-[2-(4-Methoxypyridin-2-yl)ethyl]-6-iodo-3H-imidazo[4,5-b]pyridine 608880-74-6P, 2-[2-(4-Methoxypyridin-2-yl)ethyl]-6-(4-aminophenyl)-3H-imidazo[4,5-b]pyridine RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic

preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)

(inducible NO-synthase inhibitor; preparation of alkoxypyridines as inducible NO-synthase inhibitors)

RN608880-48-4 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)ethyl]- (9CI) (CA INDEX NAME)

RN 608880-53-1 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 6-bromo-2-[2-(4-methoxy-2-pyridinyl)ethyl]-(9CI) (CA INDEX NAME)

RN608880-54-2 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 6-iodo-2-[2-(4-methoxy-2-pyridinyl)ethyl]-(9CI) (CA INDEX NAME)

608880-74-6 CAPLUS RN

CN Benzenamine, 4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5b]pyridin-6-yl]- (9CI) (CA INDEX NAME)

608880-50-8P, 2-[2-(4-Methoxypyridin-2-yl)ethyl]-7-methyl-3H-TΤ imidazo[4,5-b]pyridine 608880-51-9P, 2-[2-(4-Methoxypyridin-2yl)ethyl]-5,7-dimethyl-3H-imidazo[4,5-b]pyridine 608880-52-0P, 2-[2-(4-Methoxypyridin-2-yl)ethyl]-5-methoxy-3H-imidazo[4,5-b]pyridine 608880-55-3P, 2-[2-(4-Methoxypyridin-2-yl)ethyl]-6-nitro-3H-

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imidazo[4,5-b]pyridine 608880-56-4P, 2-[2-(4-Methoxypyridin-2-
yl)ethyl]-6-trifluoromethyl-3H-imidazo[4,5-b]pyridine 608880-57-5P
  2-[2-(4-Methoxypyridin-2-yl)ethyl]-6-phenyl-3H-imidazo[4,5-b]pyridine
608880-58-6P, 2-[2-(4-Methoxypyridin-2-yl)ethyl]-6-methyl-3H-
imidazo[4,5-b]pyridine 608880-59-7P, 2-[2-(4-Methoxypyridin-2-
y1)ethy1]-6-(2-methylpropy1)-3H-imidazo[4,5-b]pyridine
608880-60-0P, 2-[2-(4-Methoxypyridin-2-yl)ethyl]-6-
cyclohexylmethyl-3H-imidazo[4,5-b]pyridine 608880-61-1P,
2-[2-(4-Methoxypyridin-2-yl)ethyl]-6-(2-phenylethyl)-3H-imidazo[4,5-
b]pyridine 608880-62-2P, 2-[2-(4-Methoxypyridin-2-y1)ethyl]-6-
(3,4-dichlorophenyl)-3H-imidazo[4,5-b]pyridine 608880-63-3P,
2-[2-(4-Methoxypyridin-2-yl)ethyl]-6-(4-bromophenyl)-3H-imidazo[4,5-
b]pyridine 608880-64-4P, 2-[2-(4-Methoxypyridin-2-y1)ethy1]-6-(4-
bromobenzyl)-3H-imidazo[4,5-b]pyridine 608880-65-5P,
7-(2-Methoxyethoxy)-2-[2-(4-methoxypyridin-2-yl)ethyl]-3H-imidazo[4,5-
b]pyridine 608880-66-6P, 2-[2-(4-Methoxypyridin-2-yl)ethyl]-7-(2-
phenylethoxy)-3H-imidazo[4,5-b]pyridine 608880-67-7P,
2-[2-(4-Methoxypyridin-2-yl)ethyl]-7-(2,2,2-trifluoroethoxy)-3H-
imidazo[4,5-b]pyridine 608880-68-8P, 7-Hydroxy-2-[2-(4-
methoxypyridin-2-yl)ethyl]-3H-imidazo[4,5-b]pyridine 608880-69-9P
  2-[2-(4-Methoxypyridin-2-yl)ethyl]-7-(2-p-tolylethyl)-3H-imidazo[4,5-
b]pyridine 608880-70-2P, 2,7-Bis[2-(4-methoxypyridin-2-yl)ethyl]-
3H-imidazo[4,5-b]pyridine 608880-71-3P, 2-[2-(4-Methoxypyridin-2-
y1) ethyl]-7-[2-(2-pyridyl)ethyl]-3H-imidazo[4,5-b]pyridine
608880-72-4P, 2-[2-(4-Methoxypyridin-2-yl)ethyl]-6-p-tolyl-3H-
imidazo[4,5-b]pyridine 608880-73-5P, 2-[2-(4-Methoxypyridin-2-
y1)ethy1]-6-(pyridin-3-y1)-3H-imidazo[4,5-b]pyridine 608880-75-7P
, 2-[2-(4-Methoxypyridin-2-yl)ethyl]-6-(4-hydroxyphenyl)-3H-imidazo[4,5-
b]pyridine 608880-76-8P, 2-[2-(4-Methoxypyridin-2-yl)ethyl]-6-[4-
(N, N-dimethylamino)phenyl]-3H-imidazo[4,5-b]pyridine 608880-77-9P
, 2-[2-(4-Methoxypyridin-2-yl)ethyl]-6-(4-trifluoromethylphenyl)-3H-
imidazo[4,5-b]pyridine 608880-78-0P, 2-[2-(4-Methoxypyridin-2-
yl)ethyl]-6-(3,4-dimethoxyphenyl)-3H-imidazo[4,5-b]pyridine
608880-79-1P, 2-[2-(4-Methoxypyridin-2-yl)ethyl]-6-(4-
benzyloxyphenyl)-3H-imidazo[4,5-b]pyridine 608880-80-4P,
2-[2-(4-Methoxypyridin-2-yl)ethyl]-6-(4-benzyloxy-3-fluorophenyl)-3H-
imidazo[4,5-b]pyridine 608880-81-5P, 2-[2-(4-Methoxypyridin-2-
yl)ethyl]-6-(4-cyanophenyl)-3H-imidazo[4,5-b]pyridine 608880-82-6P
, 2-[2-(4-Methoxypyridin-2-yl)ethyl]-3H-imidazo[4,5-b]pyridine-6-
carboxylic acid methyl ester 608880-83-7P, N-[4-[2-[2-(4-
Methoxypyridin-2-yl)ethyl]-3H-imidazo[4,5-b]pyridin-6-yl]phenyl]acetamide
608880-84-8P, N-[4-[2-[2-(4-Methoxypyridin-2-yl)ethyl]-3H-
imidazo[4,5-b]pyridin-6-yl]phenyl]benzenesulfonamide 608880-85-9P
 2-[2-(4-Methoxy-1-oxopyridin-2-yl)ethyl]-3H-imidazo[4,5-b]pyridine
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
(Uses)
   (inducible NO-synthase inhibitor; preparation of alkoxypyridines as
   inducible NO-synthase inhibitors)
608880-50-8 CAPLUS
1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridiny1)ethy1]-7-methy1-
(9CI) (CA INDEX NAME)
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$$\begin{array}{c|c} N & N & \text{CH}_2\text{--}\text{CH}_2 \\ \hline & N & N \\ \hline & Me \end{array}$$

RN

CN

RN 608880-51-9 CAPLUS
CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)ethyl]-5,7-dimethyl(9CI) (CA INDEX NAME)

Me N N 
$$CH_2-CH_2$$
 OMe  $NH$   $NH$   $NH$ 

RN 608880-52-0 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 5-methoxy-2-[2-(4-methoxy-2-pyridinyl)ethyl]-(9CI) (CA INDEX NAME)

$$\stackrel{\text{MeO}}{\longrightarrow} \stackrel{\text{N}}{\longrightarrow} \stackrel{\text{N}}{\longrightarrow} \text{CH}_2 - \text{CH}_2 - \stackrel{\text{OMe}}{\longrightarrow}$$

RN 608880-55-3 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)ethyl]-6-nitro-(9CI) (CA INDEX NAME)

RN 608880-56-4 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)ethyl]-6-(trifluoromethyl)- (9CI) (CA INDEX NAME)

RN 608880-57-5 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridiny1)ethy1]-6-phenyl-(9CI) (CA INDEX NAME)

RN 608880-58-6 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)ethyl]-6-methyl-(9CI) (CA INDEX NAME)

$$\begin{array}{c|c} N & N & \text{CH}_2\text{-CH}_2 \\ \hline Me & NH & N \end{array}$$

RN 608880-59-7 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)ethyl]-6-(2-methylpropyl)- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ i-Bu \end{array} \qquad \begin{array}{c} N & & \\ N & & \\ N & & \\ \end{array} \qquad \begin{array}{c} OMe \\ N & \\ \end{array}$$

RN 608880-60-0 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 6-(cyclohexylmethyl)-2-[2-(4-methoxy-2-pyridinyl)ethyl]- (9CI) (CA INDEX NAME)

$$CH_2$$
 $CH_2$ 
 $CH_2$ 

RN 608880-61-1 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)ethyl]-6-(2-phenylethyl)- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ \text{Ph-} & \text{CH}_2 - \text{CH}_2 \\ \end{array} \\ \begin{array}{c} \text{OMe} \\ & \text{NH} \\ \end{array}$$

RN 608880-62-2 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 6-(3,4-dichlorophenyl)-2-[2-(4-methoxy-2-pyridinyl)ethyl]- (9CI) (CA INDEX NAME)

$$C1$$
 $H$ 
 $N$ 
 $CH_2-CH_2$ 
 $N$ 
OMe

RN 608880-63-3 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 6-(4-bromophenyl)-2-[2-(4-methoxy-2-pyridinyl)ethyl]- (9CI) (CA INDEX NAME)

RN 608880-64-4 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 6-[(4-bromophenyl)methyl]-2-[2-(4-methoxy-2-pyridinyl)ethyl]- (9CI) (CA INDEX NAME)

RN 608880-65-5 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 7-(2-methoxyethoxy)-2-[2-(4-methoxy-2-pyridinyl)ethyl]- (9CI) (CA INDEX NAME)

MeO-CH2-CH2-O

RN 608880-66-6 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)ethyl]-7-(2-phenylethoxy)- (9CI) (CA INDEX NAME)

Ph-CH2-CH2-O.

RN 608880-67-7 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)ethyl]-7-(2,2,2-trifluoroethoxy)- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} N & N & CH_2-CH_2 & OMe \\ \hline & NH & N & N \end{array}$$

F3C-CH2-0

RN 608880-68-8 CAPLUS

CN 1H-Imidazo[4,5-b]pyridin-7-ol, 2-[2-(4-methoxy-2-pyridinyl)ethyl]- (9CI) (CA INDEX NAME)

RN 608880-69-9 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)ethyl]-7-[2-(4-methylphenyl)ethyl]- (9CI) (CA INDEX NAME)

RN 608880-70-2 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2,7-bis[2-(4-methoxy-2-pyridinyl)ethyl]- (9CI) (CA INDEX NAME)

RN 608880-71-3 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)ethyl]-7-[2-(2-pyridinyl)ethyl]- (9CI) (CA INDEX NAME)

RN 608880-72-4 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)ethyl]-6-(4-methylphenyl)- (9CI) (CA INDEX NAME)

RN 608880-73-5 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)ethyl]-6-(3-pyridinyl)- (9CI) (CA INDEX NAME)

RN 608880-75-7 CAPLUS

CN Phenol, 4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]- (9CI) (CA INDEX NAME)

RN 608880-76-8 CAPLUS

CN Benzenamine, 4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]-N,N-dimethyl- (9CI) (CA INDEX NAME)

RN 608880-77-9 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)ethyl]-6-[4-(trifluoromethyl)phenyl]- (9CI) (CA INDEX NAME)

RN 608880-78-0 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 6-(3,4-dimethoxyphenyl)-2-[2-(4-methoxy-2-pyridinyl)ethyl]- (9CI) (CA INDEX NAME)

MeO 
$$H$$
  $CH_2-CH_2$   $N$   $OMe$ 

RN 608880-79-1 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)ethyl]-6-[4-(phenylmethoxy)phenyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

RN 608880-80-4 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 6-[3-fluoro-4-(phenylmethoxy)phenyl]-2-[2-(4-methoxy-2-pyridinyl)ethyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} \text{Ph-CH}_2\text{--O} & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & &$$

RN 608880-81-5 CAPLUS

CN Benzonitrile, 4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]- (9CI) (CA INDEX NAME)

RN 608880-82-6 CAPLUS

CN lH-Imidazo[4,5-b]pyridine-6-carboxylic acid, 2-[2-(4-methoxy-2-pyridinyl)ethyl]-, methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} N & N & CH_2-CH_2 & OMe \\ \hline MeO-C & NH & N & N \end{array}$$

RN 608880-83-7 CAPLUS

CN Acetamide, N-[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]- (9CI) (CA INDEX NAME)

RN 608880-84-8 CAPLUS

CN Benzenesulfonamide, N-[4-[2-[2-(4-methoxy-2-pyridinyl)ethyl]-1H-imidazo[4,5-b]pyridin-6-yl]phenyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ &$$

RN 608880-85-9 CAPLUS

CN lH-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-1-oxido-2-pyridinyl)ethyl]-(9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> d his

(FILE 'HOME' ENTERED AT 10:54:18 ON 02 MAY 2007)

FILE 'REGISTRY' ENTERED AT 10:54:30 ON 02 MAY 2007

FILE 'REGISTRY' ENTERED AT 10:55:33 ON 02 MAY 2007

STRUCTURE UPLOADED

L2 5 S L1

L3 133 S L1 FULL

FILE 'CAPLUS' ENTERED AT 10:56:11 ON 02 MAY 2007 7 S L3 FULL

=> log y

L1

L4

COST IN U.S. DOLLARS	SINCE FILE ENTRY	TOTAL SESSION
FULL ESTIMATED COST	37.83	210.59
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE ENTRY	TOTAL SESSION
CA SUBSCRIBER PRICE	-5.46	-5.46

STN INTERNATIONAL LOGOFF AT 10:57:13 ON 02 MAY 2007

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PASSWORD:

NEWS HOURS

NEWS LOGIN

NEWS IPC8

TERMINAL (ENTER 1, 2, 3, OR ?):2

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Welcome to STN International
NEWS 1
                Web Page URLs for STN Seminar Schedule - N. America
NEWS 2
        JAN 08
                CHEMLIST enhanced with New Zealand Inventory of Chemicals
NEWS 3
        JAN 16
                CA/CAplus Company Name Thesaurus enhanced and reloaded
NEWS 4
        JAN 16
                IPC version 2007.01 thesaurus available on STN
NEWS 5
        JAN 16
                WPIDS/WPINDEX/WPIX enhanced with IPC 8 reclassification data
NEWS 6 JAN 22
                CA/CAplus updated with revised CAS roles
NEWS 7
        JAN 22
                CA/CAplus enhanced with patent applications from India
NEWS 8
        JAN 29
                 PHAR reloaded with new search and display fields
NEWS 9
        JAN 29
                CAS Registry Number crossover limit increased to 300,000 in
                 multiple databases
        FEB 15
NEWS 10
                 PATDPASPC enhanced with Drug Approval numbers
        FEB 15
NEWS 11
                RUSSIAPAT enhanced with pre-1994 records
NEWS 12
        FEB 23
                KOREAPAT enhanced with IPC 8 features and functionality
NEWS 13
        FEB 26
                MEDLINE reloaded with enhancements
                EMBASE enhanced with Clinical Trial Number field
NEWS 14
        FEB 26
NEWS 15
        FEB 26
                TOXCENTER enhanced with reloaded MEDLINE
                IFICDB/IFIPAT/IFIUDB reloaded with enhancements
NEWS 16
        FEB 26
NEWS 17
        FEB 26
                CAS Registry Number crossover limit increased from 10,000
                 to 300,000 in multiple databases
NEWS 18
        MAR 15
                WPIDS/WPIX enhanced with new FRAGHITSTR display format
NEWS 19
        MAR 16
                CASREACT coverage extended
NEWS 20
        MAR 20
                MARPAT now updated daily
NEWS 21
                LWPI reloaded
        MAR 22
NEWS 22
        MAR 30
                RDISCLOSURE reloaded with enhancements
NEWS 23
        APR 02
                JICST-EPLUS removed from database clusters and STN
NEWS 24
        APR 30
                GENBANK reloaded and enhanced with Genome Project ID field
NEWS 25
        APR 30
                CHEMCATS enhanced with 1.2 million new records
NEWS 26
        APR 30
                CA/CAplus enhanced with 1870-1889 U.S. patent records
NEWS 27
        APR 30
                 INPADOC replaced by INPADOCDB on STN
        MAY 01
                New CAS web site launched
NEWS 28
NEWS EXPRESS NOVEMBER 10 CURRENT WINDOWS VERSION IS V8.01c, CURRENT
              MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),
              AND CURRENT DISCOVER FILE IS DATED 25 SEPTEMBER 2006.
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=> file reg
COST IN U.S. DOLLARS

SINCE FILE TOTAL
ENTRY SESSION
0.21 0.21

FULL ESTIMATED COST

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STRUCTURE FILE UPDATES: 1 MAY 2007 HIGHEST RN 934050-43-8 DICTIONARY FILE UPDATES: 1 MAY 2007 HIGHEST RN 934050-43-8

New CAS Information Use Policies, enter HELP USAGETERMS for details.

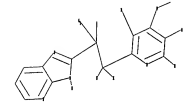
TSCA INFORMATION NOW CURRENT THROUGH December 2, 2006

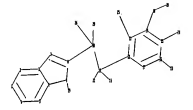
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=> Uploading C:\Program Files\Stnexp\Queries\10573203a.str





```
chain nodes :
10 11 18 19 21 22 23 24 25 26 28 29
ring nodes :
1 2 3 4 5 6 7 8 9 12 13 14 15 16 17
chain bonds :
8-10 \quad 9-19 \quad 10-11 \quad 10-26 \quad 10-29 \quad 11-12 \quad 11-21 \quad 11-22 \quad 13-25 \quad 14-18 \quad 15-24 \quad 16-23
ring bonds :
1-2 1-6 2-3 3-4 3-7 4-5 4-9 5-6 7-8 8-9 12-13 12-17 13-14 14-15 15-16
16-17
exact/norm bonds :
3-7 4-9 7-8 8-9 10-26 14-18 18-28
exact bonds :
8-10 \quad 9-19 \quad 10-11 \quad 10-29 \quad 11-12 \quad 11-21 \quad 11-22 \quad 13-25 \quad 15-24 \quad 16-23
normalized bonds :
1-2 1-6 2-3 3-4 4-5 5-6 12-13 12-17 13-14 14-15 15-16 16-17
isolated ring systems :
containing 1 : 12 :
```

## G1:C,H

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:CLASS 11:CLASS 12:Atom 13:Atom 14:Atom 15:Atom 16:Atom 17:Atom 18:CLASS 19:CLASS 21:CLASS 22:CLASS 23:CLASS 24:CLASS 25:CLASS 26:CLASS 28:CLASS 29:CLASS

=> d 11

L1 HAS NO ANSWERS

STR

Structure attributes must be viewed using STN Express query preparation.

=> s 11

SAMPLE SEARCH INITIATED 10:49:36 FILE 'REGISTRY'

SAMPLE SCREEN SEARCH COMPLETED - 13 TO ITERATE

100.0% PROCESSED

13 ITERATIONS

0 ANSWERS

4 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*

> BATCH \*\*COMPLETE\*\*

PROJECTED ITERATIONS: 44 TO 476 0 TO

PROJECTED ANSWERS:

L2 0 SEA SSS SAM L1

=> s 11 full

FULL SEARCH INITIATED 10:49:40 FILE 'REGISTRY'

FULL SCREEN SEARCH COMPLETED - 256 TO ITERATE

100.0% PROCESSED 256 ITERATIONS

SEARCH TIME: 00.00.01

4 SEA SSS FUL L1

=> file caplus

COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION

FULL ESTIMATED COST 172.10 172.31

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=> s 13 full

L4 1 L3

=> d ibib abs hitstr tot

L4 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2005:300444 CAPLUS

DOCUMENT NUMBER: 142:373835

TITLE: Preparation of imidazopyridine derivatives as

inducible NO-synthase inhibitors

INVENTOR(S): Boer, Rainer; Ulrich, Wolf-Ruediger; Eltze, Manfrid;

Marx, Degenhard; Graedler, Ulrich; Fuchss, Thomas

PATENT ASSIGNEE(S): Altana Pharma A.-G., Germany

SOURCE: PCT Int. Appl., 31 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.					KIND DATE			APPLICATION NO.						DATE					
WO	0 2005030768			A1 20050407			WO 2004-EP52370					20040930							
	W:						ΑU,												
		CN,	co,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,		
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	KZ,	LC,		
							LV,												
							PL,												
							TZ,												
	RW:						MW,												
							RU,												
							GR,												
					BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	NE,		
SN, TD, TG																			
	2004276012							AU 2004-276012											
						CA 2004-2540239													
EΡ				20060705 EP 2004-787257															
	R:						ES,									MC,	PT,		
							RO,										SK,	HR	
	1856492					2006								20040930					
	R 2004015038 A																		
JP	JP 2007507463				T		20070329 JP 2006-53					5302	60	0 20040930					

NO 2006001343 20060324 NO 2006-1343 Α 20060324 US 2007010549 Α1 20070111 US 2006-573203 20060324 PRIORITY APPLN. INFO.: EP 2003-22042 20031001 WO 2004-EP52370 W 20040930

OTHER SOURCE(S):

MARPAT 142:373835

GΙ

RN

Title compds. I [R1 = H, alkyl; R2 = H, halo, OH, etc.; R3 = H, halo, alkyl, alkoxy; R4 = alkyl; R5 = alkyl] and their resp. pharmaceutically acceptable salts, are prepared and disclosed as inducible no-synthase inhibitors. Thus, e.g., II was prepared via Wittig reaction of triphenyl-{1-(3H-imidazo[4,5-b]pyridin-2-yl)-propyl}-phosphonium chloride (preparation given) with 4-methoxypyridine-2-carboxaldehyde and subsequent hydrogenation. The activity of II towards inducible NO-synthase was evaluated in an inhibition assay and revealed a -logIC50 value of 7.15 mol/L. I as inducible NO-synthase inhibitors should prove useful in the treatment of acute and chronic inflammatory diseases.

IT 849346-44-7P 849346-45-8P 849346-46-9P

849346-44-7P 849346-45-8P 849346-46-9P RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of imidazopyridine derivs. as inducible NO-synthase inhibitors) 849346-44-7 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)-1-methylethyl]-, hydrochloride (9CI) (CA INDEX NAME)

## •x HCl

RN 849346-45-8 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[1-[(4-methoxy-2-pyridinyl)methyl]propyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} N & N & Et \\ \hline & CH-CH_2 & \\ \hline & N & N \end{array}$$
 OMe

RN 849346-46-9 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)-1,1-dimethylethyl](9CI) (CA INDEX NAME)

IT 849346-55-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of imidazopyridine derivs. as inducible NO-synthase inhibitors)

RN 849346-55-0 CAPLUS

CN 1H-Imidazo[4,5-b]pyridine, 2-[2-(4-methoxy-2-pyridinyl)-1-methylethyl]-(9CI) (CA INDEX NAME)

$$\begin{array}{c|c} N & Me \\ \hline CH-CH_2 & OMe \\ \hline NH & N & N \\ \end{array}$$

REFERENCE COUNT:

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> d his

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3

FILE 'REGISTRY' ENTERED AT 10:49:08 ON 02 MAY 2007

L1 STRUCTURE UPLOADED

L2 0 S L1

L3 4 S L1 FULL

FILE 'CAPLUS' ENTERED AT 10:49:47 ON 02 MAY 2007

L4 1 S L3 FULL

=> log y

COST IN U.S. DOLLARS

FULL ESTIMATED COST

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)

SINCE FILE

TOTAL

SESSION

5.74

178.05

CA SUBSCRIBER PRICE ENTRY SESSION
-0.78 -0.78

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TERMINAL (ENTER 1, 2, 3, OR ?):2

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NEWS
                 Web Page URLs for STN Seminar Schedule - N. America
     1
NEWS
         JAN 08
                 CHEMLIST enhanced with New Zealand Inventory of Chemicals
                 CA/CAplus Company Name Thesaurus enhanced and reloaded
NEWS
     3
         JAN 16
NEWS
     4
         JAN 16
                 IPC version 2007.01 thesaurus available on STN
NEWS
     5
         JAN 16
                 WPIDS/WPINDEX/WPIX enhanced with IPC 8 reclassification data
NEWS
         JAN 22
                 CA/CAplus updated with revised CAS roles
NEWS
         JAN 22
                 CA/CAplus enhanced with patent applications from India
NEWS
     8
         JAN 29
                 PHAR reloaded with new search and display fields
NEWS 9
         JAN 29
                 CAS Registry Number crossover limit increased to 300,000 in
                 multiple databases
NEWS 10 FEB 15
                 PATDPASPC enhanced with Drug Approval numbers
NEWS 11
        FEB 15
                 RUSSIAPAT enhanced with pre-1994 records
NEWS 12
        FEB 23
                 KOREAPAT enhanced with IPC 8 features and functionality
NEWS 13 FEB 26
                 MEDLINE reloaded with enhancements
NEWS 14 FEB 26
                 EMBASE enhanced with Clinical Trial Number field
NEWS 15 FEB 26
                 TOXCENTER enhanced with reloaded MEDLINE
NEWS 16 FEB 26
                 IFICDB/IFIPAT/IFIUDB reloaded with enhancements
NEWS 17
         FEB 26
                 CAS Registry Number crossover limit increased from 10,000
                 to 300,000 in multiple databases
NEWS 18
         MAR 15
                 WPIDS/WPIX enhanced with new FRAGHITSTR display format
NEWS 19
        MAR 16
                 CASREACT coverage extended
                 MARPAT now updated daily
NEWS 20
        MAR 20
NEWS 21
        MAR 22
                 LWPI reloaded
NEWS 22 MAR 30
                 RDISCLOSURE reloaded with enhancements
NEWS 23 APR 02
                 JICST-EPLUS removed from database clusters and STN
NEWS 24 APR 30
                 GENBANK reloaded and enhanced with Genome Project ID field
NEWS 25 APR 30
                 CHEMCATS enhanced with 1.2 million new records
NEWS 26 APR 30
                 CA/CAplus enhanced with 1870-1889 U.S. patent records
NEWS 27
         APR 30
                 INPADOC replaced by INPADOCDB on STN
NEWS 28
        MAY 01
                 New CAS web site launched
NEWS EXPRESS NOVEMBER 10 CURRENT WINDOWS VERSION IS V8.01c, CURRENT
              MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),
              AND CURRENT DISCOVER FILE IS DATED 25 SEPTEMBER 2006.
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              STN Operating Hours Plus Help Desk Availability
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              Welcome Banner and News Items
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=> file caplus

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SINCE FILE TOTAL ENTRY SESSION 0.21 0.21

FULL ESTIMATED COST

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http://www.cas.org/infopolicy.html

=> s inducible NO-synthase activity

64838 INDUCIBLE

3522859 NO

193306 NOS

1926 NOES

3634642 NO

(NO OR NOS OR NOES)

103884 SYNTHASE

5999 SYNTHASES

104972 SYNTHASE

(SYNTHASE OR SYNTHASES)

2220096 ACTIVITY

440620 ACTIVITIES

2405089 ACTIVITY

(ACTIVITY OR ACTIVITIES)

L1 71 INDUCIBLE NO-SYNTHASE ACTIVITY
(INDUCIBLE (W) NO (W) SYNTHASE (W) ACTIVITY)

=> s l1 and inflammator?

177863 INFLAMMATOR?

L2 15 L1 AND INFLAMMATOR?

=> s 12 and py<2003

22885215 PY<2003

L3 13 L2 AND PY<2003

=> d ibib abs hitstr tot

L3 ANSWER 1 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2002:344598 CAPLUS

DOCUMENT NUMBER:

137:362706

TITLE:

Effects of inhaled nitric oxide in a mouse model of sepsis-induced acute lung injury

AUTHOR(S): Razavi, Habib M.; Werhun, Robert; Scott, Jeremy A.;

Weicker, Sean; Wang, Le Feng; McCormack, David G.;

Mehta, Sanjay

CORPORATE SOURCE: A.C. Burton Vascular Research Laboratory, University

of Western Ontario, London, ON, Can.

SOURCE: Critical Care Medicine (2002), 30(4),

868-873

CODEN: CCMDC7; ISSN: 0090-3493 Lippincott Williams & Wilkins

DOCUMENT TYPE: Journal LANGUAGE: English

PUBLISHER:

Although inhaled NO transiently improves oxygenation in patients with acute lung injury, it has not affected clin. outcomes. As well, the effects of inhaled NO on the pathophysiol. features of acute lung injury were not well defined. Therefore, the authors assessed the effects of inhaled NO on the degree of pulmonary inflammation and injury in a mouse model of sepsis-induced acute lung injury. Design: Randomized, controlled animal study. Setting: Research laboratory of an academic institution. Subjects: Male C57BI/ $\acute{6}$  mice. Interventions: Sepsis was induced by cecal ligation and perforation. At the time of surgery, septic and naive mice were randomized to exposure to either 40 ppm inhaled NO or room air for 24 h before they were killed. Measurements and main results: Sepsis-induced acute lung injury was characterized by increased pulmonary myeloperoxidase (68 vs. 13 mU/mg protein in naive mice), pulmonary 8-isoprostane content (627 vs. 88 pg/mg protein in naive mice), and protein in bronchoalveolar lavage fluid. Inhaled NO exposure in septic mice completely abrogated the septic increases in myeloperoxidase activity and pulmonary 8-isoprostane content but had no effect on bronchoalveolar lavage protein. The induction of sepsis also was associated with an increase in pulmonary inducible NO synthase activity (2.8

vs. 0.4 pmolemin-lemg-1 protein in naive mice), and inhaled NO attenuated this increase in pulmonary inducible NO synthase activity. Conclusions: Exposure to inhaled NO early in the course of sepsis-induced acute lung injury is associated with reduced pulmonary leukocyte infiltration and less oxidative injury. Decreased lung inflammation and injury with inhaled NO is associated with decreased pulmonary inducible NO synthase

activity. Therefore, inhaled NO may have greater clin. benefit if administered earlier in the natural history of acute lung injury in patients.

REFERENCE COUNT: 47 THERE ARE 47 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 2 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2001:339872 CAPLUS

DOCUMENT NUMBER: 135:282996

TITLE: In vivo microvascular actions of Artemisia vulgaris L.

in a model of ischemia-reperfusion injury in the rat

intestinal mesentery

AUTHOR(S): Tigno, Xenia T.; Gumila, Elinor

CORPORATE SOURCE: Department of Physiology, College of Medicine,

University of the Philippines Manila, Manila,

Philippines

SOURCE: Clinical Hemorheology and Microcirculation (

2000), 23(2-4), 159-165

CODEN: CHMIFQ; ISSN: 1386-0291

PUBLISHER: IOS Press DOCUMENT TYPE: Journal LANGUAGE: English

AB Fractions of aqueous exts. of leaves from A. vulgaris L. (commonly known as mugwort) were tested for their effects on tissue damage brought about by ischemia-reperfusion injury in the rat mesentery. After a midline abdominal incision, the mesenteric area was exteriorized and observed by videomicroscopy. After basal observations of systemic blood pressure, heart rate, venular diams. and leukocyte adhesion along the venules, the

mesenteric artery and vein were occluded for 10 min. Prior to occlusion, treated animals were given a bolus injection of a 1% solution of a hexane-soluble fraction of the aqueous exts., while the control group received saline. Monastral Blue dye was also administered before the occlusion via the jugular vein to assess transendothelial leakage. Hemodynamic and cellular parameters were measured immediately after the release of occlusion and at 10-min intervals thereafter. The exts. had no significant effects on mean blood pressures and heart rates, but appeared to reduce leukocyte adherence and transendothelial leakage while improving flow in the ischemia-reperfused organ. The extract fractions are known to contain yomogin, which has been previously shown to inhibit inducible NO synthase activity,

which may explain the anti-inflammatory property of the plant.

REFERENCE COUNT: 27 THERE ARE 27 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 3 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2000:531501 CAPLUS

DOCUMENT NUMBER: 134:125374

TITLE: Prostaglandins and nitric oxide as molecular targets

for anti-inflammatory therapy

AUTHOR(S): Sautebin, L.

CORPORATE SOURCE: Department of Experimental Pharmacology, University of

Naples Federico II, Naples, 80131, Italy Fitoterapia (2000), 71(Suppl. 1), S48-S57

CODEN: FTRPAE; ISSN: 0367-326X

PUBLISHER: Elsevier Science B.V. DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

SOURCE:

AΒ A review with 62 refs. Nonsteroidal anti-inflammatory drugs are among the most widely used drugs worldwide, in spite of their renal and gastric side effects. Medicinal plants may represent a useful source of new and effective therapeutic agents, particularly considering new findings concerning the mediators of inflammation, such as prostaglandins The discovery of two isoforms of cyclooxygenase, which catalyzes the conversion of arachidonic acid to prostaglandins, has opened new perspectives in the treatment of inflammatory diseases. Like cyclooxygenase, NO synthase, the enzyme which converts L-arginine to NO, also exists in two isoforms. It appears that the constitutive isoforms of both enzymes (cyclooxygenase-1 and constitutive NO synthase) have a regulatory-physiol. role, whereas the inducible isoforms (cyclooxygenase-2 and inducible NO synthase) are involved in inflammation. A number of medicinal plants have been screened for their ability to inhibit cyclooxygenase-2 and/or inducible NO synthase activity and/or expression.

REFERENCE COUNT: 62 THERE ARE 62 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 4 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2000:182139 CAPLUS

DOCUMENT NUMBER: 132:319972

CORPORATE SOURCE:

TITLE: Inducible NO synthase

activity in blood vessels and heart: New insight into cell origin and consequences

AUTHOR(S): Muller, B.; Kleschyov, A. L.; Gyorgy, K.; Stoclet,

J.-C.
Pharmacologie et Physico-Chimie des Interactions

Cellulaires et, Universite Louis Pasteur de

Strasbourg, Illkirch, Fr.
SOURCE: Physiological Research (Prague) (2000),

49(1), 19-26

CODEN: PHRSEJ; ISSN: 0862-8408

PUBLISHER: Institute of Physiology, Academy of Sciences of the

Czech Republic

DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

A review, with 33 refs. Induction of the inducible form of nitric oxide synthase (iNOS) in the vascular and cardiac tissue by several inflammatory stimuli may result in the production of large amts. of nitric oxide (NO) for a sustained period. Recent data obtained in the rat aorta in which iNOS was induced by lipopolysaccharide (LPS) have demonstrated that adventitial cells represent the main site of NO production Adventitial-derived NO can exert an immediate down-regulatory effect on smooth muscle contraction (via activation of the cGMP pathway) but may also initiate longer lasting effects through the formation of NO stores within the medial layer. One candidate for such NO stores are dinitrosyl non-heme iron complexes. Low mol. weight thiols interact with preformed NO stores and promote vasorelaxation by a cGMP-independent mechanism involving the activation of potassium channels. In the heart, the induction of iNOS is involved in delayed protection against ischemia-reperfusion-induced functional damages. Recent data obtained with monophosphoryl lipid A, a non-toxin derivative of LPS, strongly suggest that iNOS-derived NO in the rat heart does not act as an immediate mediator of the cardioprotection but rather as a trigger of long-term protective mechanisms. Thus, the present data reveal the important role of adventitial cells as a site of iNOS expression and activity in intact blood vessels. The induction of adaptive mechanisms in the heart and the formation of releasable NO stores in blood vessels are examples of long-term consequences of iNOS induction. These new information are relevant for a better understanding of the circumstances in which NO overprodn. by iNOS may play either a beneficial or deleterious role in these tissues.

REFERENCE COUNT:

33 THERE ARE 33 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 5 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1999:779892 CAPLUS

DOCUMENT NUMBER:

132:73378

TITLE:

Effects of natural products on the inhibition of

lipopolysaccharide-inducible nitric oxide synthase

activity in RAW264.7 cell culture system

AUTHOR(S):

Park, Bong-Joo; Cho, Myung-Haing; Kim, Kyeong-Ho; Lee,

Sang Kook; Lee, Chong-Soon; An, Gil-Hwan; Mar,

Woongchon

CORPORATE SOURCE:

Natural Products Research Institute, Seoul National

University, Seoul, 110-460, S. Korea

SOURCE:

Natural Product Sciences (1999), 5(3),

113-120

CODEN: NPSCFB; ISSN: 1226-3907 Korean Society of Pharmacognosy

DOCUMENT TYPE:

PUBLISHER:

LANGUAGE:

Journal English

Nitric oxide (NO) is a free radical synthesized from L-arginine by nitric oxide synthase (NOS). It is believed that NO is an important mediator in numerous physiol. and inflammatory responses. Particularly, a large amount of NO released from the inducible nitric oxide synthase (iNOS) is mostly associated with inflammatory processes. Overprodn. of NO in these processes including sepsis and autoimmune diseases can have deleterious consequences and pathophysiol. relevance. Therefore, for the discovery of new inhibitory agents against iNOS activity, we have evaluated about 100 kinds of natural products after partition into three layers (n-hexane, Et acetate and aqueous) from 100% methanol exts. to study inhibitory effects on iNOS activity induced by lipopolysaccharide (LPS) in RAW264.7 cells culture system. As a pos. control, curcumin, which is known as an anti-tumor promoter, anti-inflammatory agent as an iNOS inhibitor, was used and showed the dose-dependent inhibitory effect (IC50, 2.5 μg/mL). Among tested fractions, the n-hexane fraction of Cimicifuga heracleifolia (IC50: 9.65 µg/mL), Forsythiae fructus (IC50: 6.36 μg/mL), Saposhnikovia divaricata (IC50: 5.92 μg/mL), and the Et acetate fraction of Chrysanthemum sibiricum (IC50: 2.56 µg/mL),

Gastrodia elata (IC50: 3.46  $\mu$ g/mL), and the aqueous fraction of Dianthus chinensis (IC50: 6.73  $\mu$ g/mL), Euonymus alatus (IC50: 6.78  $\mu$ g/mL), and Mechania urticifoloria (IC50: 8.01  $\mu$ g/mL) showed strong inhibitory activity against LPS-stimulated iNOS. Especially, the Et acetate fraction of Chrysanthemum sibiricum (IC50: 2.56  $\mu$ g/mL), which exhibited the strongest inhibition against iNOS, was fractionated with silica-gel column chromatog. These subfractions exhibited dose-dependent inhibition against iNOS activity in the range of 2.59-5.6  $\mu$ g/mL except for fraction Number 3, 4, 5, 6, 8, 9, and 16. Our study shows that Chrysanthemum sibiricum has the strongest inhibitory effect against iNOS activity and has similar effect to curcumin. Therefore, further studies for the identification of active principles from Chrysanthemum sibiricum and investigation for the mechanism of the inhibition of iNOS by active principles will be performed.

REFERENCE COUNT:

THERE ARE 43 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 6 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1998:475912 CAPLUS

DOCUMENT NUMBER: 129:201968

TITLE: Effects of murine recombinant interleukin-10 on the

inflammatory disease of rats transgenic for

HLA-B27 and human β2-microglobulin

AUTHOR(S): Bertrand, Viviane; Quere, Sylvie; Guimbaud, Rosine;

Sogni, Philippe; Chauvelot-Moachon, Laurence; Tulliez, Micheline; Lamarque, Dominique; Charreire, Jeannine; Giroud, Jean-Paul; Couturier, Daniel; Chaussade,

Stanislas; Breban, Maxime

CORPORATE SOURCE: Groupe de Recherche en Pathologie Digestive, Hopital

Cochin, Paris, Fr.

SOURCE: European Cytokine Network (1998), 9(2),

161-170

CODEN: ECYNEJ; ISSN: 1148-5493

PUBLISHER: John Libbey Eurotext

DOCUMENT TYPE: Journal LANGUAGE: English

Rats transgenic for  $HL\bar{A}-B27$  and human  $\beta2-microglobulin$  develop a AB spontaneous, multisystem, inflammatory disease that resembles human B27-associated disease and that involves the gut mucosa. This model predominantly affects the colon and is characterized by an extensive infiltration of the mucosa by inflammatory cells, largely composed of mononuclear cells. In addition, an increased plasma level of nitric oxide (NO)-derived metabolites was described in this model. Deficiency in the anti-inflammatory cytokine, interleukin-10 (IL-10), leads to the development of colitis in IL-10 knockout mice, suggesting that IL-10 plays a major role in the control of gut inflammation. The objectives of this work were to study the mechanisms of the inflammatory bowel disease (IBD) in HLA-B27 rats and to determine the effects of treatment with IL-10. The 33-3 line of HLA-B27 recombinant rats with established disease was treated in two consecutive expts. with murine recombinant IL-10 for five weeks. Assessment of the effect of this treatment was performed, based on clin., histol. and biol. (myeloperoxidase and inducible NO synthase activities; tumor necrosis factor- $\alpha$ , interferon- $\gamma$ , CD3, iNOS and  $\beta$ -actin mRNA expression). In 33-3 rats with established disease, mesenteric lymph nodes were hyperplastic, and colonic cellularity and MPO and iNOS activities in the colonic mucosa were increased without any detectable effects of IL-10 administration. IFN-  $\!\gamma$  and iNOS mRNA were only detected in the colon of transgenic rats. Despite a lack of effect on disease expression, IL-10 strikingly reduced the level of IFN- $\gamma$  mRNA in gut mucosa. Up-regulation of IFN- $\gamma$  mRNA suggests that the IBD of HLA-B27 rats is mediated by T-helper 1 lymphocytes. Sustained administration of IL-10, in HLA-B27 rats with established disease, efficiently inhibited IFN- $\gamma$  mRNA expression but did not influence disease expression: these results

indicate that IFN-y may exert a critical role at an earlier stage of the disease rather in the maintenance of the lesions.

REFERENCE COUNT:

THERE ARE 60 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 7 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

60

ACCESSION NUMBER: 1998:252574 CAPLUS

DOCUMENT NUMBER: 129:23093

TITLE: Tetracycline inhibits the nitric oxide synthase

activity induced by endotoxin in cultured murine

macrophages

AUTHOR(S): D'Agostino, Pietro; Arcoleo, Francesco; Barbera,

Caterina; Di Bella, Gloria; La Rosa, Marzia; Misiano,

Gabriella; Milano, Salvatore; Brai, Melchiorre;

Cammarata, Giuseppe; Feo, Salvatore; Cillari, Enrico Institute of General Pathology, University of Palermo,

CORPORATE SOURCE:

Palermo, 90134, Italy

European Journal of Pharmacology (1998), SOURCE:

346(2/3), 283-290

CODEN: EJPHAZ; ISSN: 0014-2999

PUBLISHER: Elsevier Science B.V.

DOCUMENT TYPE: Journal LANGUAGE: English

Here we investigate the effects of tetracycline base and of a semi-synthetic tetracycline derivative, doxycycline, on the induction of inducible nitric oxide synthase and, hence, on the production of nitric oxide (NO) by lipopolysaccharide in J774 macrophage cultured in vitro. treatment of J774 line with tetracycline base (6.25-250 µM) or

doxycycline (5-50  $\mu M$ ) dose-dependently decreased the

lipopolysaccharide-stimulated (1 µg/mL) inducible NO synthase activity and, consequently, nitrite formation.

For instance, the inhibition was 70% for tetracycline base at 250  $\mu M$ 

and 68% for doxycycline at 50 µM. The inhibitory effect of

tetracyclines was due neither to a reduction in the viability of the cells,

studied as colorimetric 3-[4,5-dimethylthiazol-2yl]-2,5-

diphenyltetrazolium bromide (MTT) reduction assay, nor to an indiscriminate inhibition of total protein synthesis, but to a specific decrease in inducible NO synthase protein content in the cells, as attested by the significant reduction of the expression of inducible NO synthase, assayed by sodium-dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) and Western blot. However, no effect of tetracyclines on inducible NO synthase mRNA accumulation could be demonstrated in lipopolysaccharidestimulated macrophage line, suggesting that the inhibitory effect of tetracyclines on NO synthesis involves post-transcriptional events. reduction in lipopolysaccharide-stimulated nitrite accumulation produced by tetracyclines was significantly less when they were applied 6 h after lipopolysaccharide and absent 12 h after lipopolysaccharide, indicating that tetracyclines modify an early event in inducible NO synthase activation operating after mRNA transcription. The findings presented in this study indicate that the modulation of NO synthesis is another possible pathway by which tetracyclines may function as anti-

inflammatory compds.

REFERENCE COUNT: 39 THERE ARE 39 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 8 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

1997:646568 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 127:326156

TITLE: Time-dependent actions of nitric oxide synthase

inhibition on colonic inflammation induced by

trinitrobenzene sulfonic acid in rats

AUTHOR(S): Kiss, Jozsef; Lamarque, Dominique; Delchier, Jean Charles; Whittle, Brendan J. R.

CORPORATE SOURCE: CHU Henri Mondor, INSERM U99, 51 Avenue du Marechal de

Lattre, Creteil, 94010, Fr.

SOURCE: European Journal of Pharmacology (1997),

336(2/3), 219-224

CODEN: EJPHAZ; ISSN: 0014-2999

PUBLISHER: Elsevier DOCUMENT TYPE: Journal LANGUAGE: English

The time-dependent actions following pretreatment or delayed AΒ administration of the nitric oxide (NO) synthase inhibitor,

NG-nitro-L-arginine Me ester (L-NAME) on colonic inflammation and

inducible NO synthase activity

following the intrarectal administration of trinitrobenzene sulfonic acid (TNBS) were evaluated in the rat. Intracolonic instillation of TNBS (30 mg in 0.25 mL of 50% ethanol) led to macroscopic injury, an increase of mucosal myeloperoxidase activity and the expression of the

Ca2+-independent inducible NO synthase over 8 days. The

inflammatory response following TNBS reached maximum levels between 12 and 72 h and then it declined until 14 days. Oral administration of L-NAME (25 mg kg-1 per 24 h in the drinking water) 2 days before TNBS augmented macroscopic damage and increased colonic inducible

NO synthase activity 6, 12, 24 and 72 h after

TNBS administration. In contrast, when L-NAME was administered 6 h after TNBS instillation, at time of expression of inducible NO synthase, the macroscopic lesions were reduced, as well as the enhanced inducible NO synthase activity,

determined, over 72 h. Delayed (6 h after TNBS) administration of L-NAME also attenuated the colonic myeloperoxidase activity provoked by TNBS, after 24 This activity was not affected by pretreatment (2 days before TNBS) with L-NAME. These findings indicate that the timing of administration of non-selective NO synthase inhibitors such as L-NAME, in models of colitis is critical to the eventual outcome. Thus, pretreatment with L-NAME, which will inhibit constitutive NO synthase, exacerbates the subsequent damage following challenge. In contrast, delayed administration of L-NAME at the time of inducible NO synthase expression, has a beneficial action on the colonic injury and inflammation. The findings also suggest that during the development of chronic colonic inflammation, a local overprodn. of nitric oxide by the inducible NO synthase in inflamed tissue is involved in the injury.

REFERENCE COUNT:

35 THERE ARE 35 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 9 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1996:733453 CAPLUS

DOCUMENT NUMBER: 126:98991

TITLE: 2-Amino-4-methylpyridine as a potent inhibitor of

inducible NO synthase

activity in vitro and in vivo

Faraci, W. Stephen; Nagel, Arthur A.; Verdries, AUTHOR(S):

Kimberly A.; Vincent, Lawrence A.; Xu, Hong; Nichols,

Lois E.; Labasi, Jeffrey M.; Salter, Eben D.;

Pettipher, E. Roy

CORPORATE SOURCE: Central Research Division, Pfizer Inc., Groton, CT,

06340, USA

British Journal of Pharmacology (1996), SOURCE:

119(6), 1101-1108

CODEN: BJPCBM; ISSN: 0007-1188

PUBLISHER: Stockton DOCUMENT TYPE: Journal LANGUAGE: English

The ability of 2-amino-4-methylpyridine to inhibit the catalytic activity of the inducible NO synthase (NOS II) enzyme was characterized in vitro and in vivo. In vitro, 2-amino-4-methylpyridine inhibited NOS II activity derived from mouse RAW 264.7 cells with an IC50 of 6 nM. Enzyme kinetic studies indicated that inhibition is competitive with respect to arginine. 2-Amino-4-methylpyridine was less potent on human recombinant NOS II (IC50=40 nM) and was still less potent on human recombinant NOS I and NOS

III (IC50=100 nM). NG-monomethyl-L-arginine (L-NMMA), N6-iminoethyl-L-lysine (L-NIL) and aminoquanidine were much weaker inhibitors of murine NOS II than 2-amino-4-methylpyridine but, unlike 2-amino-4-methylpyridine, retained similar activity on human recombinant NOS II. L-NMMA inhibited all three NOS isoforms with similar potency (IC50s 3-7  $\mu M$ ). In contrast, compared to activity on human recombinant NOS III, L-NIL displayed 10 + selectivity for murine NOS II and 11 + selectivity for human recombinant NOS II while aminoquanidine displayed 7.3+ selectivity for murine NOS II and 3.7+ selectivity for human recombinant NOS II. Mouse RAW 264.7 macrophages produced high levels of nitrite when cultured overnight in the presence of lipopolysaccharide (LPS) and interferon- $\gamma$ . Addition of 2-amino-4-methylpyridine at the same time as the LPS and IFN- $\gamma$ , dose-dependently reduced the levels of nitrite (IC50=1.5  $\mu$ M) without affecting the induction of NOS II protein. Increasing the extracellular concentration of arginine decreased the potency of 2-amino-4-methylpyridine but at concns. up to 10  $\mu$ M, 2-amino-4-methylpyridine did not inhibit the uptake of [3H]-arginine into the cell. Addition of 2-amino-4-methylpyridine after the enzyme was induced also dose-dependently inhibited nitrite production Together, these data suggest that 2-amino-4-methylpyridine reduces cellular production of NO by competitive inhibition of the catalytic activity of NOS II, in agreement with results obtained from in vitro enzyme kinetic studies. When infused i.v. in conscious unrestrained rats, 2-amino-4-methylpyridine inhibited the rise in plasma nitrate produced in response to i.p. injection of LPS (ID50-0.009 mg kg-1 min-1). Larger doses of 2-amino-4-methylpyridine were required to raise mean arterial pressure in untreated conscious rats (ED50=0.060 mg kg-1 min-1) indicating 6.9 + selectivity for NOS II over NOS III in vivo. Under the same conditions, L-NMMA was nonselective while L-NIL and aminoquanidine displayed 5.2 + and 8.6 + selectivity, resp. All of these compds. caused significant increases in mean arterial pressure at doses above the ID50 for inhibition of NOS II activity in vivo. 2-Amino-4-methylpyridine also inhibited LPS-induced elevation in plasma nitrate after either s.c. (ID50=0.3 mg kg-1) or oral (ID50=20.8 mg kg-1) administration. These data indicate that 2-amino-4-methylpyridine is a potent inhibitor of NOS II activity in vitro and in vivo with a similar degree of isoenzyme selectivity to that of L-NIL and aminoquanidine in rodents.

L3ANSWER 10 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1996:595100 CAPLUS

DOCUMENT NUMBER: 125:292514

S-Substituted isothioureas are potent inhibitors of TITLE:

nitric oxide biosynthesis in cartilage

AUTHOR(S):

Jang, Daniel; Szabo, Csaba; Murrell, George A. C. CORPORATE SOURCE: Laboratory for Soft Tissue Research, The Hospital for

Special Surgery, Cornell University Medical College,

New York, NY, USA

SOURCE: European Journal of Pharmacology (1996),

312(3), 341-347

CODEN: EJPHAZ; ISSN: 0014-2999

PUBLISHER: Elsevier DOCUMENT TYPE: Journal LANGUAGE: English

Nitric oxide (NO) is a multifunctional messenger mol. generated by a AB family of enzymes, the nitric oxide synthases, and is overproduced in osteoarthritis and rheumatoid arthritis. Chondrocytes are the major native source of NO in diarthrodial joints. Chondrocytic inducible nitric oxide synthase induced by inflammatory cytokines and bacterial cell wall fragments mediates many of the catabolic events in arthritis. Agents which specifically inhibit chondrocyte inducible NO synthase, may thus have a role in the management in arthritis. We evaluated a novel class of potential inducible NO synthase inhibitors, the S-substituted isothioureas, for their ability to inhibit inducible NO synthase activity in cultured bovine chondrocytes and

explants of cartilage from patients with osteoarthritis. Two isothioureas, S-Me isothiourea and S-(aminoethyl) isothiourea were 2-4 times more potent than N G-monomethyl-L-arginine monoacetate, 5-10 times more potent than aminoguanidine and over 300 times more potent than  $N\omega$ -nitro-L-arginine and  $N\omega$ -nitro-L-arginine Me ester. The rank order of potency of the NO synthase inhibitors was S-(aminoethyl) isothiourea>S-Me isothiourea>NG-monomethyl-L-arginine>aminoquanidine>N.ome ga.-nitro-L-arginine =  $N \omega$ -nitro-L-arginine Me ester. The order of potency was reversed (N $\omega$ -nitro-L-arginine Me ester =  $N\omega$ -nitro-L-arginine >NG-monomethyl-L-arginine = S-Me isothiourea>S-(aminoethyl) isothiourea>aminoguanidine) when evaluating the same compds. ability to inhibit constitutive NO synthase activity in bovine endothelial cells. In comparison to conventional arginine-based analogs, the isothioureas represent a more potent and relatively specific class of inhibitors of inducible NO synthase in cartilage and thus may be beneficial in the management of arthritis.

L3 ANSWER 11 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1996:442012 CAPLUS

DOCUMENT NUMBER: 125:132054

TITLE: CD23-mediated nitric oxide synthase pathway induction

in human keratinocytes is inhibited by retinoic acid

derivatives

AUTHOR(S): Becherel, Pierre-Andre; Le Goff, Liliane; Ktorza,

Sandra; Chosidow, Olivier; Frances, Camille; Issaly, Francoise; Mencia-Huerta, Jean-Michel; Debre, Patrice;

Mossalayi, M. Djavad; Arock, Michel

CORPORATE SOURCE: Molecular Immuno-Hematology Group, Pitie-Salpetriere

Hospital, Paris, 75013, Fr.

SOURCE: Journal of Investigative Dermatology (1996),

106(6), 1182-1186

CODEN: JIDEAE; ISSN: 0022-202X

PUBLISHER: Blackwell DOCUMENT TYPE: Journal LANGUAGE: English

AΒ Retinoids exert various functions including anti-proliferative and antiinflammatory effects on many cell types including keratinocytes and are widely used in skin diseases, such as psoriasis and acne. previously shown that human keratinocytes express low affinity IqE receptor (FceRII/CD23) when stimulated with interleukin-4. IgE ligates CD23 and induces the production of nitrites (reflecting the mobilization of the nitric oxide [NO]-pathway) and tumor necrosis factor- $\alpha$  by human keratinocytes. Here, 13-cis and all-trans retinoic acid (RA) were shown to reduce the production of nitrites by IgE-activated keratinocytes by 80% in a time- and concentration-dependent fashion. As a consequence, RA derivs. also reduced the production of tumor necrosis factor- $\alpha$  by these cells by 70%. The level of inducible NO synthase activity in activated human keratinocytes was significantly decreased upon treatment of the cells with RA derivs. (inhibition by 60% of the mean inducible NO synthase activity with 13-cis RA, 2  $\mu\text{M}$ ). Treatment for 24 h with RA derivs. almost completely abolished transcription of inducible NO synthase-specific mRNA in activated keratinocytes. Therefore, RA derivs. downregulate tumor necrosis factor- $\alpha$  release and the NO-transduction pathway through the inhibition of inducible NO synthase transcription. Together, our data provide evidence for inhibition of the NO-pathway by 13-cis and all-trans retinoic acid on CD23-activated human keratinocytes. These data may clarify the mechanism of the anti-inflammatory activity of RA derivs. in skin diseases.

ANSWER 12 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1996:149619 CAPLUS

DOCUMENT NUMBER: 124:257595

TITLE: Role of nitric oxide in induction of

. inflammatory fluid secretion by the mucosa of

the feline gallbladder

AUTHOR(S): Nilsson, Bengt; Delbro, Dick; Hedin, Lars; Conradi,

Nils; Thune, Anders; Friman, Styrbjoern; Wennmalm,

Ake; Yan, Zhong-Qun; Svanvik, Joar

CORPORATE SOURCE: Department Surgery, Sahlgrenska University Hospital,

Goteborg, Swed.

SOURCE: Gastroenterology (1996), 110(2), 598-606

CODEN: GASTAB; ISSN: 0016-5085

PUBLISHER: Saunders DOCUMENT TYPE: Journal LANGUAGE: English

Nitric oxide is synthesized from L-arginine and is metabolized to nitrate and nitrite. This study evaluates the effects of a pharmacol. blockade of NO synthesis on fluid transport by the inflamed gallbladder mucosa. Expts. were performed in cats with cholecystitis and in control animals. NO synthase activity was measured in gallbladder tissue; the enzyme was characterized by immunoblotting techniques and localized by immunofluorescence. Fluid transport and release of nitrate and nitrite by the gallbladder mucosa and bile and bile salt secretion from the liver were registered simultaneously in vivo. Fluid secretion in inflamed gallbladders was reversed to a net absorption in response to the NO synthase blockers No-nitro-L-arginine and aminoguanidine, and formation of nitrate was reduced. The effects were reversed by L-arginine. Increased levels of inducible NO synthase in inflamed gallbladders were shown by immunoblotting, by immunofluorescence (mainly in macrophages), and by Ca2+-independent [3H]citrulline formation from [3H]arginine. The NO synthase blockers had no effect on gallbladder fluid transport in normal gallbladders. Thus, increased levels of inducible NO synthase activity are shown in inflamed gallbladders, and a pharmacol. blockade of this enzyme blocks fluid secretion and decreases nitrate release from the mucosa.

ANSWER 13 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1994:506469 CAPLUS

DOCUMENT NUMBER: 121:106469

TITLE: Differential regulation of constitutive and inducible

nitric oxide production by inflammatory

stimuli murine endothelial cells

Walter, R.; Schaffner, A.; Schoedon, G. AUTHOR(S):

CORPORATE SOURCE: Dep. Med., Univ. Hosp. Zurich, Switz.

SOURCE: Biochemical and Biophysical Research Communications (

1994), 202(1), 450-5

CODEN: BBRCA9; ISSN: 0006-291X

DOCUMENT TYPE: Journal LANGUAGE: English

The murine vascular endothelial cell line send1 expresses both constitutive and inducible nitric oxide (NO) synthases. Interferon-gamma (IFNy) or endotoxin (LPS) alone inhibited constitutive NO production in a dose and time dependent manner. Addition of L-arginine had no influence on the decrease of NO production caused by IFN $\gamma$  or LPS. On the other hand, IFN $\gamma$  and LPS synergized in the induction of high output NO production Successive incubations with IFNy and LPS in different sequences revealed IFN $\gamma$  as the time setting signal for the induction of NO synthesis. These results demonstrate that LPS and IFN $\gamma$  have a direct suppressive effect on constitutive NO synthase while at the same time they strongly activate inducible NO production Thus inflammatory stimuli trigger murine vascular endothelial cells to switch from constitutive to inducible NO synthase activity.

FULL ESTIMATED COST 50.26 50.47

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L171 S INDUCIBLE NO-SYNTHASE ACTIVITY

L215 S L1 AND INFLAMMATOR?

L3 13 S L2 AND PY<2003

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